

Teprotumumab-trbw (Tepezza®)

Place of Service

Office Administration

Infusion Center Administration

Home Infusion Administration

Outpatient Facility Infusion Administration*

[*Prior authorization required – see section (1)]

HCPSC: J3241 per 10 mg

Condition listed in policy (see criteria for details)

- [Thyroid eye disease](#)

AHFS therapeutic class: Antineoplastic and Immunomodulating Agents

Mechanism of action: Insulin-like growth factor receptor (IGF-R) inhibitor

(1) Special Instructions and pertinent Information

Covered under the medical benefit, please submit clinical information for prior authorization review via fax.

(2) Prior Authorization/Medical Review is required for the following condition(s)

All requests for teprotumumab-trbw (Tepezza®) must be sent for clinical review and receive authorization prior to drug administration or claim payment.

Thyroid eye disease

1. Prescribed by or in consultation with an endocrinologist or ophthalmologist, **AND**
2. Documentation that patient's thyroxine and free triiodothyronine levels are less than 50% above or below normal limits

Covered Dose

Up to 10 mg/kg for first IV infusion, followed by 20 mg/kg IV every 3 weeks for 7 additional infusions (total treatment course = 8 infusions)

Coverage Period

Initial authorization: One treatment course (8 infusions over approximately six months)

Reauthorization if meets below: One treatment course (8 infusions over approximately six months)

1. Prescribed by or in consultation with an endocrinologist or ophthalmologist, **AND**
2. Documentation that patient's thyroxine and free triiodothyronine levels are less than 50% above or below normal limits, **AND**
3. One of the following:
 - a. Patient experienced an inadequate response to first treatment course with Tepezza (proptosis reduction of <2 mm), or
 - b. Patient experienced a relapse following treatment with Tepezza (e.g. increase in proptosis increase in clinical activity score [CAS])

ICD-10:

E05.00 [Thyrotoxicosis with diffuse goiter without thyrotoxic crisis or storm]

(3) The following condition(s) DO NOT require Prior Authorization/Preservice

PHP Medi-Cal

Teprotumumab-trbw (Tepezza®)

All requests for teprotumumab-trbw (Tepezza®) must be sent for clinical review and receive authorization prior to drug administration or claim payment.

(4) This Medication is NOT medically necessary for the following condition(s)

Coverage for a Non-FDA approved indication, requires that criteria outlined in Health and Safety Code § 1367.21, including objective evidence of efficacy and safety are met for the proposed indication.

Please refer to the Provider Manual and User Guide for more information.

(5) Additional Information

How supplied:

- 500 mg lyophilized powder (single-dose vial)

(6) References

- AHFS®. Available by subscription at <http://www.lexi.com>
- Burch HB, Perros P, Bednarczuk T, et al. Management of thyroid eye disease: a Consensus Statement by the American Thyroid Association and the European Thyroid Association. Eur Thyroid J 2022; 11: e220189.
- Douglas RS, Kahaly GJ, Ugradar S, et al. Teprotumumab efficacy, safety, and durability in longer-duration thyroid eye disease and re-treatment: OPTIC-X study. Ophthalmology 2022; 129:438-449.
- DrugDex®. Available by subscription at <http://www.micromedexsolutions.com/home/dispatch>
- Tepezza® (teprotumumab-trbw) [Prescribing information]. Deerfield, IL: Horizon Therapeutics USA, Inc.; 10/2021.

(7) Policy Update

Date of last review: 3Q2023

Date of next review: 3Q2024

Changes from previous policy version:

- Section (2): Thyroid eye disease -
 - Removed management for active, progressive disease
Rationale: In April 2023, FDA expanded the indication of Tepezza for use in thyroid eye disease regardless of disease activity or duration
 - Add coverage for retreatment
Rationale: 2022 American Thyroid Association/European Thyroid Association Consensus Statement Recommendations

*BSC Drug Coverage Criteria to Determine Medical Necessity
Reviewed by P&T Committee*